## WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



### INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5:

A61K 9/08, 9/48, 9/66
A61K 47/14, 31/10

(11) International Publication Number: WO 92/10996

(43) International Publication Date: 9 July 1992 (09.07.92)

(21) International Application Number: PCT/US91/08565 (74) Agents: SAYLES, Michael, J. et al.; Marion Merrell Dow Inc., 2110 East Galbraith Road, P.O. Box 156300, Cincinnati, OH 45215-6300 (US).

(30) Priority data: 629,540 18 December 1990 (18.12.90) US

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cinnati, OH 45215-6300 (US).

(81) Designated States: AT (European patent), AU, BE (Euro-

(81) Designated States: AT (European patent), AU, BE (European patent), CA, CH (European patent), DE (European patent), DK (European patent), ES (European patent), FI, FR (European patent), GB (European patent), GR (European patent), HU, IT (European patent), JP, KR, LU (European patent), NL (European patent), NO, SE (European patent).

**Published** 

With international search report.

(54) Title: ENHANCED BIOAVAILABILITY PHARMACEUTICAL COMPOSITION CONTAINING PROBUCOL

#### (57) Abstract

A pharmaceutical composition of probucol that both enhances bioavailability of the drug and reduces plasma drug level variability in a patient population comprising a therapeutically effective amount of probucol dissolved in a propylene glycol ester of fatty acids wherein the fatty acids are selected from the group consisting of the fatty acids represented by  $C_xH_{2x}O_2$ , wherein x is 4, 6, 8, 10, 12, 14, 16.

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## ENHANCED BIOAVAILABILITY PHARMACEUTICAL COMPOSITION CONTAINING PROBUCOL

## BACKGROUND OF THE INVENTION

- Probucol, a serum cholesterol lowering agent, is presently marketed as Lorelco® 250 and 500 mg tablets. The bioavailability of probucol from tablet dosage form is estimated to be 2-8 percent [J.F. Heeg and H. Tachizawa, Nouv. Presse Med., 9, 2990-2994 (1980)]. This poor
- 10 bioavailability is most likely caused by the extremely hydrophobic nature of probucol. Several approaches for improving the bioavailability of poorly water soluble drugs have been reported in the literature. Drugs are absorbed from the gastrointestinal tract most rapidly when adminis-
- from an oil solution may be enhanced, however, if the oil is digestible. Therefore, it was considered appropriate to develop an oil solution formulation of probucol filled in a hard gelatin capsule as one of the approaches to improving
- 20 its bioavailability. In developing such a formulation, it was unexpectedly discovered that one such formulation that increased bioavailability also resulted in reduced variability of plasma drug levels of probucol in a patient population to which the formulation was administered.

# SUMMARY OF THE INVENTION

Three new pharmaceutical dosage forms of probucol were prepared and the relative bioavailability was evaluated in One of these dosage form, a solution of probucol in Captex® 200 filled in hard gelatin capsules, was found (by 5 extrapolation) to be approximately equal to the Lorelco® 500 tablet at about 1/6 the dose. The solubility of probucol was determined in several natural and derived vegetable Captex® 200, a vegetable oil containing propylene glycol esters of caprylic (C8) and capric (C10) fatty acids, 10 provided the highest solubility for probucol and was therefore selected as the preferred vehicle for an improved probucol formulation. Also, unexpectedly, there was significantly less variability in probucol plasma drug levels with the Captex® 200 formulation. In view of the 15 increased bioavailability of probucol when administered in the Captex® 200 formulation and in light of the unexpected reduced variability in probucol plasma levels, this formulation is the subject of this application.

# DETAILED DESCRIPTION OF THE INVENTION

Probucol is a compound according to Formula I, namely 2,2'-bis (3,5-di-tertiarybutyl-4-hydroxyphenylthio)propane.

30 FORMULA I

The compounds of Formula I can be prepared as described in U.S. Patents Nos. 3,576,883, 3,786,100, 3,862,332, 3,987,500 and 4,900,757, incorporated herein by reference. More specifically, 2,2'-bis (3,5-di-tertiarybutyl-4-hydroxy phenylthio)propane can be prepared as described in U.S. Patent No. 3,576,883, also incorporated herein by reference. Alternately, this compound can be prepared according to the method set forth in U.S. Patent Nos. 4,734,527 (Kraus) or 4,861,443 (Van Effen), incorporated herein by reference. The indication for probucol is primary hypercholesterolemia.

10 Recent studies in animals have indicated that probucol has a beneficial effect on atherosclerosis independent of cholesterol lowering.

The present invention is directed towards pharmaceutical 15 compositions of probucol dissolved in a propylene glycol ester of fatty acids wherein the fatty acids are selected from the group consisting of the fatty acids represented by  $C_xH_{2x}O_2$ , wherein x is 4, 6, 8, 10, 12, 14, 16. This group specifically includes butyric acid, caproic acid, caprylic 20 acid, capric acid, lauric acid, myristic acid and palmitic acid. The most preferred embodiment of the invention is a propylene glycol ester of capric and caprylic acids, known as propylene glycol dicaprylate/dicaprate. Captex® 200 is a specific trade name for propylene glycol dicaprylate/-25 dicaprate and is supplied by Karlshamns Lipid Specialties USA, P.O. Box 569, Columbus, OH 43216-0569. Reference to Captex® 200 should not be construed as limiting and it will be understood that any reference to Captex® 200 should be construed to generically include all propylene glycol 30 dicaprylates/dicaprates. Propylene glycol dicaprylate/dicaprate is also known as Neobee 20, supplied by Stepan Co., PVO Dept., 100 W. Hunter Ave., Maywood, NJ 07607, and as Miglyol 840 , supplied by Huls America, P.O. Box 456, Piscataway, NJ 08855-0456. Captex® 300 and Capmul MCM is 35 also supplied by Karlshamns Lipid Specialties USA.

## SOLUBILITY DETERMINATION

The solubility of probucol was determined in olive, sunflower, peanut, vegetable, corn, Captex® 200 and 300, and Capmul® MCM oils. Captex® 200 is a propylene glycol ester of caprylic (C8) and capric (C10) fatty acids obtained by fractionation of certain coconut oil fatty acids and is known generically as propylene glycol dicaprylate/dicaprate. Captex® 300 is a caprylic and capric acid triglyceride obtained by fractionation and subsequent esterification of coconut oil and is known generically as caprylic/capric triglyceride. Capmul® MCM is a mono and diglyceride of caprylic and capric acids.

Eight grams of each oil were transferred into a glass

15 tube with a teflon liner screw cap, and 2.5 g of probucol
were added to each tube. The tubes were capped and shaken
by hand until the probucol particles were wetted. The tubes
were then rotated for at least 48 hours on a test tube
rotating apparatus. The solubility of probucol was

20 determined using a high performance liquid chromatography
(HPLC) assay procedure.

The solubility values (%w/v) of probucol in various oils are shown in Table I.

SOLUBILITY OF PROBUCOL IN VARIOUS OILS

Oil	Solubility (% w/v)	Oil	Solubility (% w/v)
Peanut	5.6	Corn	5.8
Olive	5.5	Captex® 200	18.2
Sunflower	5.8	Captex® 300	12.5
Safflower	5.9	Capmul® MCM	6.3
Vegetable	5.8		

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The highest solubility was observed in Captex® 200 oil. Considering a 50 mg probucol dose, and the constraints on the capsule size and its fill volume, the Captex® 200 was selected for further study. Although coconut oil is known 5 to increase the serum cholesterol level, the literature on Captex® oil indicated that the medium chain fatty acids present in this oil do not contribute to the increase in the cholesterol level. Additionally, there is also evidence that these acids are absorbed through the portal system 10 [V.K. Babayan, Lipids, 22, 417-420 (1987)] and may actually lower the cholesterol level [J.W. Stewart, K.D. Wiggers, N.L. Jacobson, P.J. Berger, Journal of Nutrition, 108, 561-566 (1978) and D. Kritchevsky, S.A. Tepper, Journal of Nutrition, 86, 67-72 (1965)]. Although the exact mechanism 15 of action of probucol is not completely understood, there is speculation that its primary mechanism of action is in the The portal absorption of the fatty acids present in Captex® 200 may be an advantage if probucol is to exercise its action mainly in the liver.

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In order to determine if the solubility of probucol could be enhanced by incorporating absolute ethanol in the oil, three binary systems: safflower oil:ethanol (90:10), polyethylene glycol (PEG) 400:ethanol (90:10, 80:20, 70:30), and Captex® 200:ethanol (95:5, 90:10, 85:15, 80:20, and 75:25) were evaluated. The solubility data shown in Table II indicate that in each case the solubility of probucol increased in the presence of ethanol.

TABLE II SOLUBILITY OF PROBUCOL IN VARIOUS BINARY SOLVENT SYSTEMS

Solvent System	Ratio	Solubility (%w/v)
Safflower Oil:Ethanol	90:10	11.5
PEG 400	100:00	2.8
PEG 400:Ethanol	90:10	5.8
PEG 400:Ethanol	80:20	9.5
PEG 400:Ethanol	70:30	13.1
Captex <sup>®</sup> 200:Ethanol	95:5	23.0
Captex <sup>®</sup> 200:Ethanol	90:10	24.0
Captex <sup>®</sup> 200:Ethanol	85:15	25.0
Captex <sup>®</sup> 200:Ethanol	80:20	26.0
Captex® 200:Ethanol	75:25	26.0

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# BIOAVAILABILITY AND SERUM VARIABILITY STUDIES

Further studies were conducted to assess the bioavailability of the experimental formulations of probucol. studies were conducted as open, randomized, parallel studies with twelve subjects per treatment group. Lorelco® 500 mg tablets were used as the reference formulation and compared with a Scherer soft gelatin capsule (Protocol A) and Captex® Oil Solution and PEG (polyethylene glycol) 8000 comelt (Protocol B), respectively, each containing 50 mg of 25 probucol.

The current formulation of Lorelco® is 500 mg of probucol in admixture with corn starch, ethylcellulose, glycerine, hydroxypropyl cellulose, hydroxypropyl methylcellulose 2910, iron oxide, lactose, magnesium stearate, microcrystalline cellulose, polysorbate 80, talc and titanium dioxide. PEG 8000 comelt is a mixture of probucol and polyethylene glycol 8000 (PEG 8000 is known in the art). Size two gelatin capsules were filled with 100 mg WO 92/10996 PCT/US91/08565

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of the 50:50 probucol:PEG 8000 comelt, corresponding to a 50 mg dose of probucol. The Scherer soft gel is a mixture of fill weight 310 mg consisting of 50 mg probucol, 208 mg Captex® 200, 26 mg polysorbate 80 and 26 mg Imwitor 742 (caprylic/capric glycerides-HULS America).

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Summary statistics (mean, standard deviation, and coefficient of variation) for dose corrected pharmacokinetic parameters are listed in Tables III and IV for Protocols B and A respectively.

TABLE III

PROTOCOL B - Summary Statistics For Dose

Corrected Pharmacokinetic Parameters

Mean ± S.D., (C.V.), (N = 12)

Dose Corrected Parameters	Lorelco®	Captex <sup>®</sup> Oil	PEG 8000
AUC168 (µg*hr ml-1)	162 ± 108	919 ± 127	979 ± 363
	(67%)	(14%)	(37%)
C <sub>MAX</sub> (µg/ml)	2.32 ± 1.50	13.5 ± 2.72	13.9 ± 4.23
	(65%)	(20%)	(30%)
T <sub>MAX</sub> (hr)	20.0 ± 7.24	18.3 ± 7.13	20.0 ± 7.43
	(36%)	(39%)	(37%)

PROTOCOL A - Summary Statistics For Dose

Corrected Pharmacokinetic Parameters

Mean ± S.D., (C.V.), (N = 12)

Dose Corrected Parameters	Lorelco®	Scherer soft gel
AUC168 (µg∗hr ml-1)	276 ± 126 (47%)	1017 ± 328 (32%)
C <sub>MAX</sub> (μg/ml)	3.74 ± 1.55 (41%)	14.2 ± 3.49 (25%)
T <sub>MAX</sub> (hr)	18.7 ± 6.89 (37%)	19.5 ± 7.14 (37%)

For Protocol B, based on the dose corrected mean AUC (area under the curve) and  $C_{max}$  (maximum concentration) values, both Captex® Oil solution and PEG 8000 comelt are estimated to be 5 times or more bioavailable than the 5 Lorelco® 500 mg tablet (Table III). Tmax values are similar for all three formulations. To make a fair comparison of the variabilities of the test formulations (Captex® Oil Solution and PEG comelt), and the reference formulation (Lorelco $^{\text{@}}$  500 mg tablet), AUC and  $C_{\text{max}}$  values for Captex $^{\text{@}}$  Oil 10 Solution and PEG 8000 comelt were multiplied by 1.76 and 1.65, respectively (these scale factors were used so that all formulations have the same AUC values). Based on the standard deviation of these extrapolated values (Table V) and the coefficient of variation of the raw values (Table 15 III), AUC values of Captex® Oil Solution are approximately 25 and 7 times less variable than the Lorelco® tablet and the PEG 8000 treatment, respectively, and  $C_{\text{max}}$  values of Captex® Oil Solution are approximately 9 and 2.3 times less variable than the Lorelco® tablet and PEG 8000 treatment, 20 respectively. The variability of PEG 8000 comelt is similar to the Lorelco® 500 mg tablet.

Similar procedures were also used for Protocol A. Based on the dose corrected values, the Scherer soft gel treatment is estimated to be 3.7 times more bioavailable than the Lorelco® 500 mg tablet (Table IV). Tmax values are similar for both formulations. The variability of Scherer soft gel treatment is similar to the Lorelco® 500 mg tablet (Table V).

<u>TABLE V</u>

<u>Comparison Of Variances With Matched AUC Values</u>

Standard Deviation Of The Extrapolated Values (N=12)

		PROTOCOL B		
5	***************************************	Treatment'	<u> </u>	
	<u>Parameter</u>	Lorelco®	PEG 8000	Captex® Oil
	AUC168 (µg*hr ml-1)	108.2	61.8	22.9
10	CMAX (µg/ml)	1.50	0.72	0.49
	TMAX (hr)	7.20	7.10	7.40

#### PROTOCOL A 15 Treatment\* Lorelco® Scherer soft Parameter 125.7 86.9 AUC168 ( $\mu g * hr ml^{-1}$ ) 0.94 1.55 20 CMAX (µg/ml) 6.90 7.10 TMAX (hr)

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### EXAMPLE I

Probucol (50.0 mg) was dissolved in propylene glycol esters of caprylic/capric fatty acids [Captex® 200, manufactured and supplied by Karlshamns Lipid Specialties USA, P.O. Box 569, Columbus, OH 43216-0569, as Captex® 200, (283.0 mg) and stirred until a clear solution was obtained. The resulting clear solution was filled into hard gelatin capsules (white opaque gelatin capsule size no. 1, 73 mg) so that each capsule contained an approximate weight of 333.0

<sup>\*</sup>Treatments with a common bracket are not significantly different.

mg of solution. The bulk solution was assayed for producor before filling the capsule and the fill weight was adjusted according to the actual percent of probucol in the solution to provide a 50 mg dose of probucol. Using a capsule banding apparatus, a solution of gelatin (0.647 mg), polysorbate 80 (0.027 mg), and purified water (2.076 mg) was applied to seal the cap to the body of the capsule. The gelatin band was then allowed to harden.

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## WHAT IS CLAIMED IS:

1. A pharmaceutical composition of probucol adapted to enhance the bioavailability of probucol while reducing plasma probucol level variability in a patient population comprising a therapeutically effective amount of probucol dissolved in a propylene glycol ester of fatty acids wherein the fatty acids are selected from the group consisting of the fatty acids represented by C<sub>x</sub>H<sub>2x</sub>O<sub>2</sub>, wherein x is 4, 6, 8, 10, 12, 14, 16.

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2. A pharmaceutical composition according to claim 1 wherein the fatty acids are selected from the group consisting of the fatty acids represented by  $C_xH_{2x}O_2$ , wherein x is 6, 8, 10, 12.

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- 3. A pharmaceutical composition according to claim 1 wherein the fatty acids are selected from the group consisting of capric and caprylic acids.
- 20 4. Use of a pharmaceutical composition according to any of claims 1-3 to lower serum cholesterol levels.

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	CT MATTER (if several classification		
According to International Patent Int.Cl.5 A 61 K 47/14	Classification (IPC) or to both National A 61 K 9/08 A A 61 K 31/10	Classification and IPC 61 K 9/48 A 61 K 9/	/66
II. FIELDS SEARCHED			
	Minimum Docur	mentation Searched?	
Classification System		Classification Symbols	
Int.C1.5	A 61 K		·
	Documentation Searched othe to the Extent that such Document	er than Minimum Documentation ts are Included in the Fields Searched <sup>8</sup>	
III. DOCUMENTS CONSIDER	ED TO BE RELEVANT <sup>9</sup>		
Category ° Citation of D	ocument, 11 with indication, where appro	priate, of the relevant passages 12	Relevant to Claim No.13
Januar 12: ex	3862332 (J.W. BARNHAR by 1975, see the claim kample 9; column 13, e le 17 (cited in the ap	s 1,3,8,11-12; column xample 14; column 14,	1,4
A US,A,4 Februa XVIII	4902513 (J. CARVAIS) ary 1990, see the clai	20 ms; column 4, example	1,4
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IV. CERTIFICATION	Cata Lawrendered Carach	Date of Mailing of this International Se	arch Report
Date of the Actual Completion of 22-01		1 7. 02. 92.	
International Searching Authori	ty PEAN PATENT OFFICE	Signature of Authorized Officer	anielle van der Haas

## ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

US 9108565

SA 54190

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 06/02/92

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Patent document cited in search report	Publication date	Patent family member(s)	Publication date
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